

1 We claim:

2 1. A patterned phosphor structure having red, green and blue sub-pixel phosphor elements  
3 for an AC electroluminescent display, comprising:

4 at least a first and a second phosphor, each emitting light in different ranges of the  
5 visible spectrum, but whose combined emission spectra contains red, green and blue light;

6 said at least first and second phosphors being in a layer, arranged in adjacent, repeating  
7 relationship to each other to provide a plurality of repeating at least first and second phosphor  
8 deposits; and

9 one or more means associated with one or more of the at least first and second  
10 phosphor deposits, and which together with the at least first and second phosphor deposits,  
11 form the red, green and blue sub-pixel phosphor elements, for setting and equalizing the  
12 threshold voltages of the red, green and blue sub-pixel phosphor elements, and for setting the  
13 relative luminosities of the red, green and blue sub-pixel phosphor elements so that they bear  
14 set ratios to one another at each operating modulation voltage used to generate the desired  
15 luminosities for red, green and blue.

16 2. The phosphor structure as set forth in claim 1, wherein the at least first and second  
17 phosphor deposits are formed from phosphors of different host materials.

18 3. The phosphor structure as set forth in claim 2, wherein the set luminosity ratios remain  
19 substantially constant over the range of operating modulation voltages.

20 4. The phosphor structure as set forth in claim 3, wherein the set luminosities ratios  
21 between the red, green and blue sub-pixel phosphor elements is about 3:6:1.

22 5. The phosphor structure as set forth in claim 2, 3 or 4, wherein the means for setting and  
23 equalizing the threshold voltages, and for setting the relative luminosities, comprises a  
24 threshold voltage adjustment layer of a dielectric material or a semiconductor material located  
25 in one or more of the positions of over, under and embedded within one or more of the at least  
26 first and second phosphor deposits.

27 6. The phosphor structure as set forth in claim 2, 3, 4 or 5, wherein the means for setting  
28 and equalizing the threshold voltages, and for setting the relative luminosities, comprises the at  
29 least first and second phosphor deposits being formed with different thicknesses.

30 7. The phosphor structure as set forth in claim 5 or 6, wherein, the means for setting and  
31 equalizing the threshold voltages, and for setting the relative luminosities, further comprises  
32 varying one or both of the following:

i. the areas of the phosphor deposits; and

ii. the concentrations of a dopant or co-dopant in the phosphor deposits.

8. The phosphor structure as set forth in claim 7, wherein the at least first and second phosphor deposits are formed from a zinc sulfide phosphor and a strontium sulfide phosphor.

9. The phosphor structure as set forth in claim 8, wherein the blue sub-pixel elements, and optionally the green sub-pixel elements are formed with a strontium sulfide phosphor, and wherein the red sub-pixel elements, and optionally the green sub-pixel elements are formed from one or more zinc sulfide phosphors.

10. The phosphor structure as set forth in claim 9, wherein the strontium sulfide phosphor is  $\text{SrS:Ce}$  and wherein the zinc sulfide phosphor is one or both of  $\text{ZnS:Mn}$  or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with  $x$  being between 0.1 and 0.3.

11. The phosphor structure as set forth in claim 8, wherein the first phosphor is  $\text{SrS:Ce}$  and the second phosphor is one or more of  $\text{ZnS:Mn}$  or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with  $x$  being between 0.1 and 0.3, and wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises a further layer of  $\text{SrS:Ce}$  over the first and second phosphor deposits, whereby the blue sub-pixel elements are provided by  $\text{SrS:Ce}$  and the red and green sub-pixel elements are provided by  $\text{SrS:Ce}$  and one or both of  $\text{ZnS:Mn}$  or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ .

12. The phosphor structure as set forth in claim 10, wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises a threshold voltage adjustment layer over the red and green sub-pixel phosphor deposits.

13. The phosphor structure as set forth in claim 10, 11 or 12, wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises the phosphor deposits being formed with different thicknesses.

14. The phosphor structure as set forth in claim 10, 11, 12 or 13, wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises varying the areas of one or more of the sub-pixel phosphor deposits.

15. The phosphor structure as set forth claim 1, 2, or 14, wherein the means for setting and equalizing the threshold voltages, and for setting the relative luminosities, comprises a threshold voltage adjustment layer selected from the group consisting of one or more of a dielectric material or a semiconductor material, which, at its deposited thickness, does not conduct until the voltage across the patterned phosphor structure exceeds the threshold voltage

1 which the patterned phosphor structure would have without the threshold voltage adjustment  
2 layer.

3 16. The phosphor structure as set forth in claim 15, wherein the threshold voltage  
4 adjustment layer is selected from the group consisting of binary metal oxides, binary metal  
5 sulfides, silica and silicon oxynitride.

6 17. The phosphor structure as set forth in claim 15, wherein the threshold voltage  
7 adjustment layer is selected from the group consisting of alumina, tantalum oxide, zinc sulfide,  
8 strontium sulfide, silica and silicon oxynitride.

9 18. The phosphor structure as set forth in claim 15, wherein the threshold voltage  
10 adjustment layer is selected from the group consisting of alumina and zinc sulfide.

11 19. The phosphor structure as set forth in claim 15, wherein threshold voltage adjustment  
12 layer is matched with the at least first or second phosphor deposits, such that if the phosphor  
13 deposit is formed from a zinc sulfide phosphor, the threshold voltage adjustment layer, if  
14 needed with that phosphor deposit, is a binary metal oxide.

15 20. The phosphor structure as set forth in claim 19, wherein the binary metal oxide is  
16 alumina when the phosphor deposit is one or more of  $\text{ZnS:Mn}$  or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with x being  
17 between 0.1 and 0.3.

18 21. The phosphor structure as set forth in claim 5, 6 or 7, wherein the means for setting and  
19 equalizing the threshold voltages and for setting the relative luminosities comprises an  
20 additional phosphor layer deposited in one or more of the positions of over, under and  
21 embedded within the at least first and second phosphor deposits, having a same or different  
22 composition from the at least first and second phosphor deposits.

23 22. The phosphor structure as set forth in claim 5, 6 or 7, wherein the first and second  
24 phosphor deposits are a strontium sulfide phosphor providing the blue sub-pixel elements and  
25 a zinc sulfide phosphor providing the red and green sub-pixel elements, and wherein the  
26 means for setting and equalizing the threshold voltages and for setting the relative luminosities  
27 is a threshold voltage adjustment layer selected from the group consisting of one or more of a  
28 dielectric material or a semiconductor material in one or more of the positions of over, under  
29 and embedded within the zinc sulfide phosphor deposits.

30 23. The phosphor structure as set forth in claim 22, wherein the phosphors are  $\text{SrS:Ce}$ ,  
31 which may be codoped with phosphorus, and  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with x being between 0.1 and  
32 0.3, and wherein the threshold voltage adjustment layer is a layer of alumina located over the

1  $\text{Zn}_{1-x}\text{Mg}_x\text{S}:\text{Mn}$  phosphor deposits.

2 24. The phosphor structure as set forth in claim 5, 6 or 7, wherein the first and second  
3 phosphor deposits are a strontium sulfide phosphor providing the blue sub-pixel elements and  
4 one or more layers of a zinc sulfide phosphor providing the red and green sub-pixel elements,  
5 and wherein the means for setting and equalizing the threshold voltages and for setting the  
6 relative luminosities is the strontium sulfide phosphor deposits being formed thicker and wider  
7 than the zinc sulfide phosphor deposits.

8 25. The phosphor structure as set forth in claim 24, wherein the phosphors are  $\text{SrS}:\text{Ce}$  for  
9 the blue sub-pixel elements, which may be codoped with phosphorus, and for the red and  
10 green sub-pixels,  $\text{Zn}_{1-x}\text{Mg}_x\text{S}:\text{Mn}$  between layers of  $\text{ZnS}:\text{Mn}$ , with  $x$  being between 0.1 and 0.3.

11 26. The phosphor structure as set forth in claim 5, 6 or 7, wherein the first and second  
12 phosphor deposits are a strontium sulfide phosphor providing the blue and green sub-pixel  
13 elements and a zinc sulfide phosphor providing the red sub-pixel elements, and wherein the  
14 means for setting and equalizing the threshold voltages and for setting the relative luminosities  
15 is a threshold voltage adjustment layer selected from the group consisting of one or more of a  
16 dielectric material or a semiconductor material in one or more of the position of over, under  
17 and embedded within the zinc sulfide phosphor deposits.

18 27. The phosphor structure as set forth in claim 26, wherein the phosphors are  $\text{SrS}:\text{Ce}$ ,  
19 which may be codoped with phosphorus, and  $\text{ZnS}:\text{Mn}$ , and wherein the threshold voltage  
20 adjustment layer is a layer of alumina located over the  $\text{ZnS}:\text{Mn}$  phosphor deposits.

21 28. An EL laminate for use in an AC electroluminescent display, comprising:

22 a rigid rear substrate;

23 a patterned phosphor structure comprising:

24 at least a first and a second phosphor, each emitting light in different ranges of  
25 the visible spectrum, but whose combined emission spectra contains red, green  
26 and blue light;

27 said at least first and second phosphors being in a layer, arranged in adjacent,  
28 repeating relationship to each other to provide a plurality of repeating at least  
29 first and second phosphor deposits; and

30 one or more means associated with one or more of the at least first and second  
31 phosphor deposits, and which together with the at least first and second  
32 phosphor deposits, form the red, green and blue sub-pixel phosphor elements,

1 for setting and equalizing the threshold voltages of the red, green and blue sub-  
2 pixel phosphor elements, and for setting the relative luminosities of the red,  
3 green and blue sub-pixel phosphor elements so that they bear set ratios to one  
4 another at each operating modulation voltage used to generate the desired  
5 luminosities for red, green and blue;

6 front and rear column and row electrodes on either side of the phosphor structure, the  
7 rows or columns of the front or rear electrode being aligned with the phosphor sub-pixel  
8 elements;

9 a thick film dielectric layer below the patterned phosphor structure formed from a  
10 sintered ceramic material having a dielectric constant greater than 500, and having a thickness  
11 greater than about 10  $\mu\text{m}$ ; and

12 optionally, optical colour filter means aligned with the red, green and blue phosphor  
13 sub-pixel elements for transmitting red, green and blue light emitted from the phosphor sub-  
14 pixel elements.

15 29. The EL laminate as set forth in claim 28, wherein the at least first and second phosphor  
16 deposits are formed from phosphors of different host materials.

17 30. The EL laminate as set forth in claim 29, wherein the set luminosity ratios remain  
18 substantially constant over the range of operating modulation voltages.

19 31. The EL laminate as set forth in claim 30, wherein the set luminosities ratios between  
20 the red, green and blue sub-pixel phosphor elements is about 3:6:1.

21 32. The EL laminate as set forth in claim 29, 30 or 31, wherein the means for setting and  
22 equalizing the threshold voltages, and for setting the relative luminosities, comprises a  
23 threshold voltage adjustment layer of a dielectric material or a semiconductor material located  
24 in one or more of the positions of over, under and embedded within one or more of the at least  
25 first and second phosphor deposits.

26 33. The EL laminate as set forth in claim 29, 30, 31, or 32, wherein the means for setting  
27 and equalizing the threshold voltages, and for setting the relative luminosities, comprises the at  
28 least first and second phosphor deposits being formed with different thicknesses.

29 34. The EL laminate as set forth in claim 32 or 33, wherein, the means for setting and  
30 equalizing the threshold voltages, and for setting the relative luminosities, further comprises  
31 varying one or both of the following:

32 i. the areas of the phosphor deposits; and

ii. the concentrations of a dopant or co-dopant in the phosphor deposits.

35. The EL laminate as set forth in claim 34, wherein the at least first and second phosphor deposits are formed from a zinc sulfide phosphor and a strontium sulfide phosphor.

36. The EL laminate as set forth in claim 35, wherein the blue sub-pixel elements, and optionally the green sub-pixel elements are formed with a strontium sulfide phosphor, and wherein the red sub-pixel elements, and optionally the green sub-pixel elements are formed from one or more zinc sulfide phosphors.

37. The EL laminate as set forth in claim 36, wherein the strontium sulfide phosphor is SrS:Ce and wherein the zinc sulfide phosphor is one or more of ZnS:Mn or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with x being between 0.1 and 0.3.

38. The EL laminate as set forth in claim 35, wherein the first phosphor is SrS:Ce and the second phosphor is one or more of ZnS:Mn or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with x being between 0.1 and 0.3, and wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises a further layer of SrS:Ce over the first and second phosphor deposits, whereby the blue sub-pixel elements are provided by SrS:Ce and the red and green sub-pixel elements are provided by SrS:Ce and one or both of ZnS:Mn or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ .

39. The EL laminate as set forth in claim 37, wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises a threshold voltage adjustment layer over the red and green sub-pixel phosphor deposits.

40. The EL laminate as set forth in claim 37, 38, or 39, wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises the phosphor deposits being formed with different thicknesses.

41. The EL laminate as set forth in claim 37, 38, 39 or 40, wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities comprises varying the areas of one or more of the sub-pixel phosphor deposits.

42. The EL laminate as set forth claim 28, 29, or 41, wherein the means for setting and equalizing the threshold voltages, and for setting the relative luminosities, comprises a threshold voltage adjustment layer selected from the group consisting of one or more of a dielectric material or a semiconductor material, which, at its deposited thickness, does not conduct until the voltage across the patterned phosphor structure exceeds the threshold voltage which the patterned phosphor structure would have without the threshold voltage adjustment layer.

1 43. The EL laminate as set forth in claim 42, wherein the threshold voltage adjustment  
2 layer is selected from the group consisting of binary metal oxides, binary metal sulfides, silica  
3 and silicon oxynitride.

4 44. The EL laminate as set forth in claim 42, wherein the threshold voltage adjustment  
5 layer is selected from the group consisting of alumina, tantalum oxide, zinc sulfide, strontium  
6 sulfide, silica and silicon oxynitride.

7 45. The EL laminate as set forth in claim 42, wherein the threshold voltage adjustment  
8 layer is selected from the group consisting of alumina and zinc sulfide.

9 46. The EL laminate as set forth in claim 42, wherein threshold voltage adjustment layer is  
10 matched with the at least first or second phosphor deposits, such that if the phosphor deposit is  
11 formed from a zinc sulfide phosphor, the threshold voltage adjustment layer, if needed with  
12 that phosphor deposit, is a binary metal oxide.

13 47. The EL laminate as set forth in claim 46, wherein the binary metal oxide is alumina  
14 when the phosphor deposit is one or more of  $\text{ZnS:Mn}$  or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with x being between  
15 0.1 and 0.3.

16 48. The EL laminate as set forth in claim 32, 33 or 34, wherein the means for setting and  
17 equalizing the threshold voltages and for setting the relative luminosities comprises an  
18 additional phosphor layer deposited in one or more of the positions of over, under and  
19 embedded within the at least first and second phosphor deposits, having a same or different  
20 composition from the at least first and second phosphor deposits.

21 49. The EL laminate as set forth in claim 32, 33 or 34, wherein the first and second  
22 phosphor deposits are a strontium sulfide phosphor providing the blue sub-pixel elements and  
23 a zinc sulfide phosphor providing the red and green sub-pixel elements, and wherein the  
24 means for setting and equalizing the threshold voltages and for setting the relative luminosities  
25 is a threshold voltage adjustment layer selected from the group consisting of one or more of a  
26 dielectric material or a semiconductor material in one or more of the positions of over, under  
27 and embedded within the zinc sulfide phosphor deposits.

28 50. The EL laminate as set forth in claim 49, wherein the phosphors are  $\text{SrS:Ce}$ , which  
29 may be codoped with phosphorus, and  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with x being between 0.1 and 0.3, and  
30 wherein the threshold voltage adjustment layer is a layer of alumina located over the  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$   
31 phosphor deposits.

32 51. The EL laminate as set forth in claim 32, 33 or 34, wherein the first and second

phosphor deposits are a strontium sulfide phosphor providing the blue sub-pixel elements and one or more layers of a zinc sulfide phosphor providing the red and green sub-pixel elements, and wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities is the strontium sulfide phosphor deposits being formed thicker and wider than the zinc sulfide phosphor deposits.

52. The EL laminate as set forth in claim 51, wherein the phosphors are SrS:Ce for the blue sub-pixel elements, which may be codoped with phosphorus, and for the red and green sub-pixels,  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$  between layers of ZnS:Mn, with x being between 0.1 and 0.3.

53. The EL laminate as set forth in claim 32, 33 or 34, wherein the first and second phosphor deposits are a strontium sulfide phosphor providing the blue and green sub-pixel elements and a zinc sulfide phosphor providing the red sub-pixel elements, and wherein the means for setting and equalizing the threshold voltages and for setting the relative luminosities is a threshold voltage adjustment layer selected from the group consisting of one or more of a dielectric material or a semiconductor material in one or more of the position of over, under and embedded within the zinc sulfide phosphor deposits.

54. The EL laminate as set forth in claim 53, wherein the phosphors are SrS:Ce, which may be codoped with phosphorus, and ZnS:Mn, and wherein the threshold voltage adjustment layer is a layer of alumina located over the ZnS:Mn phosphor deposits.

55. The EL laminate as set forth in claims 28, 29, 32, 33 or 34, wherein the thick film dielectric layer is formed from a pressed, sintered ceramic material having, compared to an unpressed, sintered dielectric layer of the same composition, improved dielectric strength, reduced porosity and uniform luminosity in an EL laminate.

56. The EL laminate as set forth in claim 35, 50, 52, or 54, wherein the thick film dielectric layer is formed from a pressed, sintered ceramic material having, compared to an unpressed, sintered dielectric layer of the same composition, improved dielectric strength, reduced porosity and uniform luminosity in an EL laminate.

57. The EL laminate as set forth in claim 55 or 56, wherein the dielectric layer has been pressed by cold isostatic pressing to reduce the thickness, after sintering, by about 20 to 50%.

58. The EL laminate as set forth in claim 57, wherein the pressed ceramic material has a reduced thickness, after sintering, of 30 to 40%.

59. The EL laminate as set forth in claim 58, wherein the pressed ceramic material has a thickness, after sintering, of between 10 and 50  $\mu\text{m}$ .



1 60. The EL laminate as set forth in claim 58, wherein the pressed ceramic material has a  
2 thickness, after sintering, of between 10 and 20  $\mu\text{m}$ .

3 61. The EL laminate as set forth in claim 60, wherein the ceramic material is a ferroelectric  
4 ceramic material having a dielectric constant greater than 500.

5 62. The EL laminate as set forth in claim 61, wherein the ceramic material has a perovskite  
6 crystal structure.

7 63. The EL laminate as set forth in claim 62, wherein the ceramic material is selected from  
8 the group consisting of one or more of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN and PMN-PT.

9 64. The EL laminate as set forth in claim 62, wherein the ceramic material is selected from  
10 the group consisting of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN and PMN-PT.

11 65. The EL laminate as set forth in claim 62, wherein the ceramic material is PMN-PT.

12 66. The EL laminate as set forth in claim 62, 64, or 65, wherein a second ceramic material  
13 is formed on the pressed, sintered dielectric layer to further smooth the surface.

14 67. The EL laminate as set forth in claim 59, wherein the second ceramic material is a  
15 ferroelectric ceramic material deposited by sol gel techniques followed by heating to convert to  
16 a ceramic material.

17 68. The EL laminate as set forth in claim 67, wherein the second ceramic material has a  
18 dielectric constant of at least 20 and a thickness of at least about 1  $\mu\text{m}$ .

19 69. The EL laminate as set forth in claim 68, wherein the second ceramic material has a  
20 dielectric constant of at least 100.

21 70. The EL laminate as set forth in claim 69, wherein the second ceramic material has a  
22 thickness in the range of 1 to 3  $\mu\text{m}$ .

23 71. The EL laminate as set forth in claim 70, wherein the second ceramic material is a  
24 ferroelectric ceramic material having a perovskite crystal structure.

25 72. The EL laminate as set forth in claim 71, wherein the second ceramic material is lead  
26 zirconium titanate or lead lanthanum zirconate titanate.

27 73. The EL laminate as set forth in claim 72, wherein the substrate and the rear electrode  
28 are formed from materials which can withstand temperatures of about 850°C.

29 74. The EL laminate as set forth in claim 73, wherein the substrate is an alumina sheet.

30 75. The EL laminate as set forth in claim 55, 66 or 72, which further comprises, a diffusion  
31 barrier layer above the dielectric layer or above the second ceramic material, which diffusion  
32 barrier layer is composed of a metal-containing electrically insulating binary compound that is

1 chemically compatible with any adjacent layers and which is precisely stoichiometric.

2 76. The EL laminate as set forth in claim 75, wherein the diffusion barrier layer is formed  
3 from a compound which differs from its precise stoichiometric composition by less than 0.1  
4 atomic percent.

5 77. The EL laminate as set forth in claim 76, wherein the diffusion barrier layer is formed  
6 from alumina, silica, or zinc sulfide.

7 78. The EL laminate as set forth in claim 76, wherein the diffusion barrier is formed from  
8 alumina.

9 79. The EL laminate as set forth in claim 77 or 78, wherein the diffusion barrier has a  
10 thickness of 100 to 1000 Å.

11 80. The EL laminate as set forth in claim 55, 66, 72 or 75, which further comprises, an  
12 injection layer above the dielectric layer, the second ceramic material or the barrier diffusion  
13 barrier, to provide a phosphor interface, composed of a binary, dielectric material which is  
14 non-stoichiometric in its composition and having electrons in a range of energy for injection  
15 into the phosphor layer.

16 81. The EL laminate as set forth in claim 80, wherein the injection layer is formed from a  
17 material which has greater than 0.5% atomic deviation from its stoichiometric composition.

18 82. The EL laminate as set forth in claim 81, wherein the injection layer is formed from  
19 hafnia or yttria.

20 83. The EL laminate as set forth in claim 82, wherein the injection layer has a thickness of  
21 100 to 1000 Å.

22 84. The EL laminate as set forth in claim 75 or 80, wherein an injection layer of hafnia is  
23 included with a phosphor formed from a zinc sulfide phosphor, and wherein a diffusion barrier  
24 layer of zinc sulfide is used with a phosphor formed from a strontium sulfide phosphor.

25 85. A method of forming a patterned phosphor structure having red, green and blue sub-  
26 pixel elements for an AC electroluminescent display, comprising:

27 selecting at least a first and a second phosphor, each emitting light in different ranges  
28 of the visible spectrum, but whose combined emission spectra contains red, green and blue  
29 light;

30 depositing and patterning said at least first and second phosphors in a layer to form a  
31 plurality of repeating at least first and second phosphor deposits arranged in adjacent,  
32 repeating relationship to each other; and

1 providing one or more means associated with one or more of the at least first and  
2 second phosphor deposits, and which together with the at least first and second phosphor  
3 deposits, form the red, green and blue sub-pixel phosphor elements, for setting and equalizing  
4 the threshold voltages of the red, green and blue sub-pixel phosphor elements and for setting  
5 the relative luminosities of the red, green and blue sub-pixel elements so that they bear set  
6 ratios to one another at each modulation voltage used to generate the desired luminosities for  
7 red, green and blue; and

8 optionally annealing the patterned phosphor structure so formed.

9 86. The method as set forth in claim 85, wherein the at least first and second phosphor  
10 deposits are formed from phosphors of different host materials.

11 87. The method as set forth in claim 86, wherein the set luminosity ratios remain  
12 substantially constant over the range of operating modulation voltages.

13 88. The method as set forth in claim 87, wherein the set luminosities ratios between the  
14 red, green and blue sub-pixel phosphor elements are about 3:6:1.

15 89. The method as set forth in claim 86, 87 or 88, wherein the patterning of the at least first  
16 and second phosphor is achieved by photolithographic techniques, including the steps of:

17 a) depositing a layer of a first phosphor which is to form at least one of the red, green  
18 or blue sub-pixel elements;

19 b) removing the first phosphor in regions which are to define the other of the red, green  
20 or blue sub-pixel elements, leaving spaced first phosphor deposits;

21 c) depositing the second phosphor material over the first phosphor deposits and in  
22 regions which are to define the other of the red, green and blue sub-pixel elements; and

23 d) removing the second phosphor material from above the first phosphor deposits  
24 leaving a plurality of repeating first and second phosphor deposits arranged in adjacent,  
25 repeating relationship to each other.

26 90. The method as set forth in claim 89, wherein step b) includes:

27 applying a photo-resist to the first phosphor, exposing the photo-resist through a photo-  
28 mask, developing the photo-resist, removing the first phosphor in regions that first phosphor is  
29 to define as one or more of the red, green and blue sub-pixel elements;

30 and wherein step d) includes:

31 removing by lift-off, the second phosphor and the resist from above the first phosphor  
32 deposits.

1 91. The method as set forth in claim 90, wherein the photo-resist in step b) is a negative  
2 resist that is exposed in the regions that the first phosphor is to define as one or more of the  
3 red, green and blue sub-pixel elements.

4 92. The method as set forth in claim 91, wherein the patterning is achieved with only one  
5 photo-mask.

6 93. The method as set forth in claim 86, 87, 88 or 91, wherein the means for setting and  
7 equalizing the threshold voltages, and for setting the relative luminosities, comprises a  
8 threshold voltage adjustment layer selected from the group consisting of one or more of a  
9 dielectric material or a semiconductor material deposited in one or more of the positions of  
10 over, under and embedded within one or more of the at least first and second phosphor  
11 deposits.

12 94. The method as set forth in claim 86, 87, 88, 91 or 93, wherein the means for setting  
13 and equalizing the threshold voltages, and for setting the relative luminosities, comprises the at  
14 least first and second phosphor deposits being deposited with different thicknesses.

15 95. The method as set forth in claim 93 or 94, wherein, the means for setting and  
16 equalizing the threshold voltages, and for setting the relative luminosities, further comprises  
17 varying one or both of the following:

- 18 i. the areas of the phosphor deposits; and
- 19 ii. the concentrations of a dopant or co-dopant in the phosphor deposits.

20 96. The method as set forth in claim 95, wherein the at least first and second phosphor  
21 deposits include a zinc sulfide phosphor and a strontium sulfide phosphor.

22 97. The method as set forth in claim 96, wherein the blue sub-pixel elements, and  
23 optionally the green sub-pixel elements are formed with a strontium sulfide phosphor, and  
24 wherein the red sub-pixel elements, and optionally the green sub-pixel elements are formed  
25 from one or more zinc sulfide phosphors.

26 98. The method as set forth in claim 97, wherein the strontium sulfide phosphor is SrS:Ce  
27 and wherein the zinc sulfide phosphor is one or more of ZnS:Mn or  $Zn_{1-x}Mg_xS:Mn$ , with x  
28 being between 0.1 and 0.3.

29 99. The method as set forth in claim 96, wherein the first phosphor is SrS:Ce and the  
30 second phosphor is one or more of ZnS:Mn or  $Zn_{1-x}Mg_xS:Mn$ , with x being between 0.1 and  
31 0.3, and wherein the means for setting and equalizing the threshold voltages and for setting the  
32 relative luminosities is provided by depositing a further layer of SrS:Ce over the first and

1 second phosphor deposits, whereby the blue sub-pixel elements are provided by SrS:Ce and  
2 the red and green sub-pixel elements are provided by SrS:Ce and one or more of ZnS:Mn or  
3  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ .

4 100. The method as set forth in claim 98, wherein the means for setting and equalizing the  
5 threshold voltages and for setting the relative luminosities are provided by depositing a  
6 threshold voltage adjustment layer over one or more of the red and green sub-pixel phosphor  
7 deposits.

8 101. The method as set forth in claim 98, 99 or 100, wherein the means for setting and  
9 equalizing the threshold voltages and for setting the relative luminosities is provided by  
10 depositing the phosphor, and thus forming the phosphor deposits, with different thicknesses.

11 102. The method as set forth in claim 98, 99, 100 or 101, wherein the means for setting and  
12 equalizing the threshold voltages and for setting the relative luminosities is provided by  
13 varying the areas of one or more of the sub-pixel phosphor deposits.

14 103. The method as set forth claim 85, 86 or 102, wherein the means for setting and  
15 equalizing the threshold voltages, and for setting the relative luminosities, is provided by  
16 depositing over one or more of the red, green and blue sub-pixel deposits, a threshold voltage  
17 adjustment layer selected from the group consisting of one or more of a dielectric material or a  
18 semiconductor material, which, at its deposited thickness, does not conduct until the voltage  
19 across the patterned phosphor structure exceeds the threshold voltage which the patterned  
20 phosphor structure would have without the threshold voltage adjustment layer.

21 104. The method as set forth in claim 103, wherein the threshold voltage adjustment layer is  
22 selected from the group consisting of binary metal oxides, binary metal sulfides, silica and  
23 silicon oxynitride.

24 105. The method as set forth in claim 103, wherein the threshold voltage adjustment layer is  
25 selected from the group consisting of alumina, tantalum oxide, zinc sulfide, strontium sulfide,  
26 silica and silicon oxynitride.

27 106. The method as set forth in claim 103, wherein the threshold voltage adjustment layer is  
28 selected from the group consisting of alumina and zinc sulfide.

29 107. The method as set forth in claim 103, wherein threshold voltage adjustment layer is  
30 matched with the at least first or second phosphor deposits, such that if the phosphor deposit is  
31 formed from a zinc sulfide phosphor, the threshold voltage adjustment layer, if needed with  
32 that phosphor deposit, is a binary metal oxide, and if the phosphor deposit is formed from a

1 strontium sulfide phosphor, the threshold voltage adjustment layer, if needed with that  
2 phosphor deposit, is a binary metal sulfide.

3 108. The method as set forth in claim 107, wherein the binary metal oxide is alumina when  
4 the phosphor deposit is one or more of  $\text{ZnS:Mn}$  or  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with  $x$  being between 0.1  
5 and 0.3.

6 109. The method as set forth in claim 93, 94, or 98, wherein the means for setting and  
7 equalizing the threshold voltages and for setting the relative luminosities comprises an  
8 additional phosphor layer deposited in one or more of the positions of over, under and  
9 embedded within the at least first and second phosphor deposits, having a same or different  
10 composition from the at least first and second phosphor deposits.

11 110. The method as set forth in claim 93, 94 or 95, wherein the first and second phosphor  
12 deposits are a strontium sulfide phosphor providing the blue sub-pixel elements and a zinc  
13 sulfide phosphor providing the red and green sub-pixel elements, and wherein the means for  
14 setting and equalizing the threshold voltages and for setting the relative luminosities is  
15 provided by depositing a threshold voltage adjustment layer selected from the group consisting  
16 of one or more of a dielectric material or a semiconductor material in one or more of the  
17 positions of over, under and embedded within the zinc sulfide phosphor deposits.

18 111. The method as set forth in claim 110, wherein the phosphors are  $\text{SrS:Ce}$ , which may be  
19 codoped with phosphorus, and  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , with  $x$  being between 0.1 and 0.3, and wherein  
20 the threshold voltage adjustment layer is a layer of alumina deposited over the  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$   
21 phosphor deposits.

22 112. The method as set forth in claim 93, 94 or 95, wherein the first and second phosphor  
23 deposits are a strontium sulfide phosphor providing the blue sub-pixel elements and one or  
24 more layers of a zinc sulfide phosphor providing the red and green sub-pixel elements, and  
25 wherein the means for setting and equalizing the threshold voltages and for setting the relative  
26 luminosities is provided by forming the strontium sulfide phosphor deposits thicker and wider  
27 than and the zinc sulfide phosphor deposits.

28 113. The method as set forth in claim 112, wherein the phosphors are  $\text{SrS:Ce}$  for the blue  
29 sub-pixel elements, which may be codoped with phosphorus, and for the red and green sub-  
30 pixels,  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$  between layers of  $\text{ZnS:Mn}$ , with  $x$  being between 0.1 and 0.3.

31 114. The method as set forth in claim 93, 94 or 95, wherein the first and second phosphor  
32 deposits are a strontium sulfide phosphor providing the blue and green sub-pixel elements and

1 a zinc sulfide phosphor providing the red sub-pixel elements, and wherein the means for  
2 setting and equalizing the threshold voltages and for setting the relative luminosities is  
3 provided by depositing a threshold voltage adjustment layer selected from the group consisting  
4 of one or more of a dielectric material or a semiconductor material in one or more of the  
5 positions of over, under and embedded within the zinc sulfide phosphor deposits.

6 115. The method as set forth in claim 114, wherein the phosphors are SrS:Ce, which may be  
7 codoped with phosphorus, and ZnS:Mn, and wherein the threshold voltage adjustment layer is  
8 a layer of alumina deposited over the ZnS:Mn phosphor deposits.

9 116. The method as set forth in claims 91, wherein one or both of the first and second  
10 phosphors is susceptible to hydrolysis, wherein the negative resist is a polyisoprene-based  
11 resist, wherein the first phosphor is removed with an acid etchant solution, and wherein the  
12 second phosphor is removed with a non-aqueous, predominately polar, aprotic solvent  
13 solution.

14 117. The method as set forth in claim 116, wherein the first and second phosphor deposits  
15 are a strontium sulfide phosphor and a zinc sulfide phosphor, and wherein the predominately  
16 polar, aprotic solvent solution is toluene, with a minor amount of methanol.

17 118. The method as set forth in claim 117, wherein the first and second phosphor deposits  
18 are patterned in a layer from SrS:Ce and ZnS:Mn, and an additional phosphor layer of SrS:Ce  
19 is deposited over the patterned layer such that, the SrS:Ce deposits form the blue sub-pixel  
20 elements, and the ZnS:Mn deposits overlaid with the SrS:Ce deposits form the red and green  
21 sub-pixel elements, the patterning being achieved by:

- 22 a) depositing a layer of the SrS:Ce which is to form the blue sub-pixel elements;
- 23 b) applying the negative photoresist on the SrS:Ce, exposing the photoresist in those  
24 regions which are to form the blue sub-pixel elements, and removing the SrS:Ce and the  
25 unexposed photoresist in those regions which are to define the red and green sub-pixel  
26 elements, leaving spaced SrS:Ce deposit;
- 27 c) depositing the ZnS:Mn to cover both the SrS:Ce deposits and the regions where the  
28 SrS:Ce has been removed;
- 29 d) optionally depositing an injection layer;
- 30 e) removing by lift-off, the ZnS:Mn, the photoresist and the optional injection layer in  
31 the regions above SrS:Ce, to form a plurality of repeating first and second phosphor deposits  
32 arranged in adjacent, repeating relationship to each other; and

1 f) providing the means for setting and equalizing the threshold voltages and setting the  
2 relative luminosities by depositing an additional layer of SrS:Ce over the first and second  
3 phosphor deposits.

4 119. The method as set forth in claim 117, wherein the first and second phosphor deposits  
5 are a strontium sulfide phosphor providing the blue sub-pixel elements and a zinc sulfide  
6 phosphor providing the red and green sub-pixel elements, and wherein the means for setting  
7 and equalizing the threshold voltages is a threshold voltage adjustment layer selected from the  
8 group consisting of one or more of a dielectric material or a semiconductor material deposited  
9 in one or more of the positions of over, under and embedded within the zinc sulfide phosphor  
10 deposits.

11 120. The method as set forth in claim 119, wherein the phosphors are SrS:Ce, which may be  
12 codoped with phosphorus, and  $Zn_{1-x}Mg_xS:Mn$ , with x being between 0.1 and 0.3, wherein the  
13 threshold voltage adjustment layer is a layer of alumina deposited over the  $Zn_{1-x}Mg_xS:Mn$   
14 phosphor, and wherein the patterning is achieved by:

15 a) depositing a layer of the SrS:Ce which is to form the blue sub-pixel elements;  
16 b) applying the negative photoresist on the SrS:Ce, exposing the photoresist in those  
17 regions which are to form the blue sub-pixel elements, and removing the SrS:Ce and the  
18 unexposed photoresist in those regions which are to define the red and green sub-pixel  
19 elements, leaving spaced SrS:Ce deposits;

20 c) depositing the  $Zn_{1-x}Mg_xS:Mn$  to cover both the SrS:Ce deposits and the regions  
21 where the SrS:Ce has been removed;

22 d) optionally depositing an injection layer;

23 e) depositing the threshold voltage adjustment layer above the  $Zn_{1-x}Mg_xS:Mn$ ; and

24 e) removing by lift-off, the  $Zn_{1-x}Mg_xS:Mn$ , the photoresist, the threshold voltage  
25 adjustment layer, and the optional injection layer in the regions above SrS:Ce, to form a  
26 plurality of repeating first and second phosphor deposits arranged in adjacent, repeating  
27 relationship to each other.

28 121. The method as set forth in claim 117, wherein the first and second phosphor deposits  
29 are a strontium sulfide phosphor providing the blue sub-pixel elements and a zinc sulfide  
30 phosphor providing the red and green sub-pixel elements, and wherein the means for setting  
31 and equalizing the threshold voltages and setting the relative luminosities is provided by  
32 forming the strontium sulfide phosphor deposits thicker and with greater area than the zinc



1 sulfide phosphor deposits.

2 122. The method as set forth in claim 121, wherein the phosphors are SrS:Ce, which may be  
3 codoped with phosphorus, and  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$  between layers of ZnS:Mn, with x being  
4 between 0.1 and 0.3, and wherein the patterning is achieved by:

- 5 a) depositing a layer of the SrS:Ce which is to form the blue sub-pixel elements;
- 6 b) applying the negative photoresist on the SrS:Ce, exposing the photoresist in those  
7 regions which are to form the blue sub-pixel elements, and removing the SrS:Ce and the  
8 unexposed photoresist in those regions which are to define the red and green sub-pixel  
9 elements, leaving spaced SrS:Ce deposits;
- 10 c) depositing the a layer of ZnS:Mn, then a layer of  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , and then a layer of  
11 ZnS:Mn to cover both the SrS:Ce deposits and the regions where the SrS:Ce has been  
12 removed;
- 13 d) optionally depositing an injection layer;
- 14 e) removing by lift-off, the ZnS:Mn and the  $\text{Zn}_{1-x}\text{Mg}_x\text{S:Mn}$ , the photoresist, and the  
15 optional injection layer in the regions above SrS:Ce, to form a plurality of repeating first and  
16 second phosphor deposits arranged in adjacent, repeating relationship to each other.

17 123. The method as set forth in claim 117, wherein the first and second phosphor deposits  
18 are a strontium sulfide phosphor providing the blue and green sub-pixel elements and a zinc  
19 sulfide phosphor providing the red sub-pixel elements, and wherein the means for setting and  
20 equalizing the threshold voltages is provided by depositing a threshold voltage adjustment  
21 layer selected from the group consisting of one or more of a dielectric material or a  
22 semiconductor material in one or more of the positions of over, under and embedded within  
23 the zinc sulfide phosphor deposits.

24 124. The method as set forth in claim 123, wherein the phosphors are SrS:Ce, which may be  
25 codoped with phosphorus, and ZnS:Mn, wherein the threshold voltage adjustment layer is a  
26 layer of alumina located over the ZnS:Mn phosphor, and wherein the patterning is achieved  
27 by:

- 28 a) depositing a layer of the SrS:Ce which is to form the blue and green sub-pixel  
29 elements;
- 30 b) applying the negative photoresist on the SrS:Ce, exposing the photoresist in those  
31 regions which are to form the blue and green sub-pixel elements, and removing the SrS:Ce and  
32 the unexposed photoresist in those regions which are to define the red sub-pixel elements,

1 leaving spaced SrS:Ce deposits for the blue and green sub-pixel elements which are wider than  
2 the regions left for the red sub-pixel elements;

3 c) depositing an optional layer of alumina as a barrier diffusion layer;

4 d) depositing the ZnS:Mn to cover both the SrS:Ce deposits and the regions where the  
5 SrS:Ce has been removed;

6 e) depositing the threshold voltage adjustment layer above the ZnS:Mn; and

7 f) removing by lift-off, the optional barrier diffusion layer, the ZnS:Mn, the  
8 photoresist, and the threshold voltage adjustment layer in the regions above SrS:Ce, to form a  
9 plurality of repeating first and second phosphor deposits arranged in adjacent, repeating  
10 relationship to each other.

11 125. A method of forming a thick film dielectric layer in an EL laminate of the type  
12 including one or more phosphor layers sandwiched between a front and a rear electrode, the  
13 phosphor layer being separated from the rear electrode by the thick film dielectric layer,  
14 comprising:

15 depositing a ceramic material in one or more layers by a thick film technique to form a  
16 dielectric layer having a thickness of 10 to 300  $\mu\text{m}$ ;

17 pressing the dielectric layer to form a densified layer with reduced porosity and surface  
18 roughness; and

19 sintering the dielectric layer to form a pressed, sintered dielectric layer which, in an EL  
20 laminate, has an improved uniform luminosity over an unpressed, sintered dielectric layer of  
21 the same composition.

22 126. The method as set forth in claim 125, wherein the dielectric layer is deposited on a  
23 rigid substrate providing the rear electrode.

24 127. The method as set forth in claim 125, wherein the pressing is isostatic pressing.

25 128. The method as set forth in claim 126, wherein the pressing is cold isostatic pressing at  
26 up to 350,000 kPa to reduce the thickness of the dielectric layer, after sintering, by about 20 to  
27 50%.

28 129. The method as set forth in claim 128, wherein the ceramic material is deposited by  
29 screen printing, in one or more layers, and is dried prior to pressing.

30 130. The method as set forth in claim 129, wherein the ceramic material is pressed to reduce  
31 the thickness, after sintering, by 30 to 40%.

32 131. The method as set forth in claim 130, wherein the ceramic material is pressed to a

thickness, after sintering, of between 10 and 50  $\mu\text{m}$ .

132. The method as set forth in claim 130, wherein the ceramic material is pressed to a thickness, after sintering, of between 10 and 20  $\mu\text{m}$ .

133. The method as set forth in claim 132, wherein the dielectric layer has a deposited thickness of 20 to 50  $\mu\text{m}$ .

134. The method as set forth in claim 132 or 133, wherein the ceramic material is a ferroelectric ceramic material having a dielectric constant greater than 500.

135. The method as set forth in claim 134, wherein the ceramic material has a perovskite crystal structure.

136. The method as set forth in claim 135, wherein the ceramic material is selected from the group consisting of one or more of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN and PMN-PT.

137. The method as set forth in claim 135, wherein the ceramic material is selected from the group consisting of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN and PMN-PT.

138. The method as set forth in claim 137, wherein the ceramic material is PMN-PT.

139. The method as set forth in claim 136, 137, or 138, wherein a second ceramic material is formed on the pressed, sintered dielectric layer to further smooth the surface.

140. The method as set forth in claim 139, wherein the second ceramic material is a ferroelectric ceramic material which is deposited by a sol gel technique to form a sol gel layer.

141. The method as set forth in claim 140, wherein the second ceramic material has a dielectric constant of at least 20 and a thickness of at least about 1  $\mu\text{m}$ .

142. The method as set forth in claim 141, wherein the second ceramic material has a dielectric constant of at least 100.

143. The method as set forth in claim 142, wherein the second ceramic material has a thickness in the range of 1 to 3  $\mu\text{m}$ .

144. The method as set forth in claim 143, wherein the second ceramic material is deposited by a sol gel techniques selected from spin deposition or dipping, followed by heating to convert to a ceramic material.

145. The method as set forth in claim 144, wherein the second ceramic material is a ferroelectric ceramic material having a perovskite crystal structure.

146. The method as set forth in claim 145, wherein the second ceramic material is lead zirconium titanate or lead lanthanum zirconate titanate.

147. The method as set forth in claim 125, 139 or 146, which further comprises, prior to

forming the dielectric layer, providing a substrate having sufficient rigidity to support the laminate, and forming the rear electrode on the substrate.

148. The method as set forth in claim 147, wherein the substrate and the rear electrode are formed from materials which can withstand temperatures of about 850°C.

149. The method as set forth in claim 148, wherein the substrate is an alumina sheet.

150. The method as set forth in claim 125, 139 or 149, which further comprises, depositing a diffusion barrier layer above the dielectric layer or above the second ceramic material, which diffusion barrier layer is composed of a metal-containing electrically insulating binary compound that is chemically compatible with any adjacent layers and which is precisely stoichiometric.

151. The method as set forth in claim 150, wherein the diffusion barrier layer is formed from a compound which differs from its precise stoichiometric composition by less than 0.1 atomic percent.

152. The method as set forth in claim 151, wherein the diffusion barrier layer is formed from alumina, silica, or zinc sulfide.

153. The method as set forth in claim 152, wherein the diffusion barrier is formed from alumina.

154. The method as set forth in claim 153, wherein the diffusion barrier has a thickness of 100 to 1000 Å.

155. The method as set forth in claim 125, 139 or 150, which further comprises, depositing an injection layer above the dielectric layer, the second ceramic material or the barrier diffusion barrier, to provide a phosphor interface, composed of a binary, dielectric material which is non-stoichiometric in its composition and having electrons in a range of energy for injection into the phosphor layer.

156. The method as set forth in claim 155, wherein the injection layer is formed from a material which has greater than 0.5% atomic deviation from its stoichiometric composition.

157. The method as set forth in claim 156, wherein the injection layer is formed from hafnia or yttria.

158. The method as set forth in claim 157, wherein the injection layer has a thickness of 100 to 1000 Å.

159. The method as set forth in claim 156 or 158, wherein the injection layer is hafnia when the phosphor is a zinc sulfide phosphor, and wherein a diffusion barrier layer of zinc sulfide is

1 used with a strontium sulfide phosphor.

2 160. A combined substrate and dielectric layer component for use in an EL laminate,  
3 comprising:

4 a substrate providing a rear electrode; and

5 a thick film dielectric layer formed on the substrate from a pressed, sintered ceramic  
6 material having, compared to an unpressed, sintered dielectric layer of the same composition,  
7 improved dielectric strength, reduced porosity and uniform luminosity in an EL laminate.

8 161. The combined substrate and dielectric layer component as set forth in claim 160,  
9 formed on a rigid substrate providing a rear electrode.

10 162. The combined substrate and dielectric layer component as set forth in claim 161,  
11 wherein the dielectric layer has been pressed by cold isostatic pressing to reduce the thickness,  
12 after sintering, by about 20 to 50%.

13 163. The combined substrate and dielectric layer component as set forth in claim 162,  
14 wherein the pressed ceramic material has a reduced thickness, after sintering, of 30 to 40%.

15 164. The combined substrate and dielectric layer component as set forth in claim 163,  
16 wherein the pressed ceramic material has a thickness, after sintering, of between 10 and 50  
17  $\mu\text{m}$ .

18 165. The combined substrate and dielectric layer component as set forth in claim 163,  
19 wherein the pressed ceramic material has a thickness, after sintering, of between 10 and 20  
20  $\mu\text{m}$ .

21 166. The combined substrate and dielectric layer component as set forth in claim 165,  
22 wherein the ceramic material is a ferroelectric ceramic material having a dielectric constant  
23 greater than 500.

24 167. The combined substrate and dielectric layer component as set forth in claim 166,  
25 wherein the ceramic material has a perovskite crystal structure.

26 168. The combined substrate and dielectric layer component as set forth in claim 167,  
27 wherein the ceramic material is selected from the group consisting of one or more of  $\text{BaTiO}_3$ ,  
28  $\text{PbTiO}_3$ , PMN and PMN-PT.

29 169. The combined substrate and dielectric layer component as set forth in claim 167,  
30 wherein the ceramic material is selected from the group consisting of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN  
31 and PMN-PT.

32 170. The combined substrate and dielectric layer component as set forth in claim 167,

wherein the ceramic material is PMN-PT.

171. The combined substrate and dielectric layer component as set forth in claim 168, 169, or 170, wherein a second ceramic material is formed on the pressed, sintered dielectric layer to further smooth the surface.

172. The combined substrate and dielectric layer component as set forth in claim 171, wherein the second ceramic material is a ferroelectric ceramic material deposited by sol gel techniques followed by heating to convert to a ceramic material.

173. The combined substrate and dielectric layer component as set forth in claim 172, wherein the second ceramic material has a dielectric constant of at least 20 and a thickness of at least about 1  $\mu\text{m}$ .

174. The combined substrate and dielectric layer component as set forth in claim 173, wherein the second ceramic material has a dielectric constant of at least 100.

175. The combined substrate and dielectric layer component as set forth in claim 174, wherein the second ceramic material has a thickness in the range of 1 to 3  $\mu\text{m}$ .

176. The combined substrate and dielectric layer component as set forth in claim 175, wherein the second ceramic material is a ferroelectric ceramic material having a perovskite crystal structure.

177. The combined substrate and dielectric layer component as set forth in claim 176, wherein the second ceramic material is lead zirconium titanate or lead lanthanum zirconate titanate.

178. The combined substrate and dielectric layer component as set forth in claim 160, 171, or 177, wherein the combined substrate and dielectric layer component is formed on a rigid substrate, on which is formed the rear electrode.

179. The combined substrate and dielectric layer component as set forth in claim 178, wherein the substrate and the rear electrode are formed from materials which can withstand temperatures of about 850°C.

180. The combined substrate and dielectric layer component as set forth in claim 179, wherein the substrate is an alumina sheet.

181. The combined substrate and dielectric layer component as set forth in claim 160, 171, or 178, which further comprises, a diffusion barrier layer above the dielectric layer or above the second ceramic material, which diffusion barrier layer is composed of a metal-containing electrically insulating binary compound that is chemically compatible with any adjacent layers

and which is precisely stoichiometric.

182. The combined substrate and dielectric layer component as set forth in claim 181, wherein the diffusion barrier layer is formed from a compound which differs from its precise stoichiometric composition by less than 0.1 atomic percent.

183. The combined substrate and dielectric layer component as set forth in claim 182, wherein the diffusion barrier layer is formed from alumina, silica, or zinc sulfide.

184. The combined substrate and dielectric layer component as set forth in claim 182, wherein the diffusion barrier is formed from alumina.

185. The combined substrate and dielectric layer component as set forth in claim 183 or 184, wherein the diffusion barrier has a thickness of 100 to 1000 Å.

186. The combined substrate and dielectric layer component as set forth in claim 160, 171, 178 or 181, which further comprises, an injection layer above the dielectric layer, the second ceramic material or the barrier diffusion barrier, to provide a phosphor interface, composed of a binary, dielectric material which is non-stoichiometric in its composition and having electrons in a range of energy for injection into the phosphor layer.

187. The combined substrate and dielectric layer component as set forth in claim 186, wherein the injection layer is formed from a material which has greater than 0.5% atomic deviation from its stoichiometric composition.

188. The combined substrate and dielectric layer component as set forth in claim 187, wherein the injection layer is formed from hafnia or yttria.

189. The combined substrate and dielectric layer component as set forth in claim 188, wherein the injection layer has a thickness of 100 to 1000 Å.

190. The combined substrate and dielectric layer component as set forth in claim 187 or 189, wherein the injection layer is hafnia with a zinc sulfide phosphor, and wherein a diffusion barrier layer of zinc sulfide is used with a strontium sulfide phosphor.

191. An EL laminate, comprising:

a planar phosphor layer;

a front and rear planar electrode on either side of the phosphor layer;

a rear substrate providing the rear electrode, the rear substrate having sufficient rigidity to support the laminate; and

a thick film dielectric layer on the rigid substrate providing the rear electrode, the thick film dielectric layer being formed from a pressed, sintered ceramic material having, compared

1 to an unpressed, sintered dielectric layer of the same composition, improved dielectric  
2 strength, reduced porosity and uniform luminosity in an EL laminate.

3 192. The EL laminate as set forth in claim 191, formed on a rigid substrate providing a rear  
4 electrode.

5 193. The EL laminate as set forth in claim 191 or 192, wherein the dielectric layer has been  
6 pressed by cold isostatic pressing to reduce the thickness, after sintering, by about 20 to 50%.

7 194. The EL laminate as set forth in claim 193, wherein the pressed ceramic material has a  
8 reduced thickness, after sintering, of 30 to 40%.

9 195. The EL laminate as set forth in claim 194, wherein the pressed ceramic material has a  
10 thickness, after sintering, of between 10 and 50  $\mu\text{m}$ .

11 196. The EL laminate as set forth in claim 194, wherein the pressed ceramic material has a  
12 thickness, after sintering, of between 10 and 20  $\mu\text{m}$ .

13 197. The EL laminate as set forth in claim 196, wherein the ceramic material is a  
14 ferroelectric ceramic material having a dielectric constant greater than 500.

15 198. The EL laminate as set forth in claim 197, wherein the ceramic material has a  
16 perovskite crystal structure.

17 199. The EL laminate as set forth in claim 198, wherein the ceramic material is selected  
18 from the group consisting of one or more of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN and PMN-PT.

19 200. The EL laminate as set forth in claim 198, wherein the ceramic material is selected  
20 from the group consisting of  $\text{BaTiO}_3$ ,  $\text{PbTiO}_3$ , PMN and PMN-PT.

21 201. The EL laminate as set forth in claim 198, wherein the ceramic material is PMN-PT.

22 202. The EL laminate as set forth in claim 199, 200 or 201, wherein a second ceramic  
23 material is formed on the pressed, sintered dielectric layer to further smooth the surface.

24 203. The EL laminate as set forth in claim 202, wherein the second ceramic material is a  
25 ferroelectric ceramic material deposited by sol gel techniques followed by heating to convert to  
26 a ceramic material.

27 204. The EL laminate as set forth in claim 203, wherein the second ceramic material has a  
28 dielectric constant of at least 20 and a thickness of at least about 1  $\mu\text{m}$ .

29 205. The EL laminate as set forth in claim 204, wherein the second ceramic material has a  
30 dielectric constant of at least 100.

31 206. The EL laminate as set forth in claim 205, wherein the second ceramic material has a  
32 thickness in the range of 1 to 3  $\mu\text{m}$ .



- 1 207. The EL laminate as set forth in claim 206, wherein the second ceramic material is a  
2 ferroelectric ceramic material having a perovskite crystal structure.
- 3 208. The EL laminate as set forth in claim 207, wherein the second ceramic material is lead  
4 zirconium titanate or lead lanthanum zirconate titanate.
- 5 209. The EL laminate as set forth in claim 191, 202, or 208, wherein the EL laminate is  
6 formed on a rigid substrate, on which is formed the rear electrode.
- 7 210. The EL laminate as set forth in claim 209, wherein the substrate and the rear electrode  
8 are formed from materials which can withstand temperatures of about 850°C.
- 9 211. The EL laminate as set forth in claim 210, wherein the substrate is an alumina sheet.
- 10 212. The EL laminate as set forth in claim 191, 202, or 209, which further comprises, a  
11 diffusion barrier layer above the dielectric layer or above the second ceramic material, which  
12 diffusion barrier layer is composed of a metal-containing electrically insulating binary  
13 compound that is chemically compatible with any adjacent layers and which is precisely  
14 stoichiometric.
- 15 213. The EL laminate as set forth in claim 212, wherein the diffusion barrier layer is formed  
16 from a compound which differs from its precise stoichiometric composition by less than 0.1  
17 atomic percent.
- 18 214. The EL laminate as set forth in claim 213, wherein the diffusion barrier layer is formed  
19 from alumina, silica, or zinc sulfide.
- 20 215. The EL laminate as set forth in claim 213, wherein the diffusion barrier is formed from  
21 alumina.
- 22 216. The EL laminate as set forth in claim 214 or 215, wherein the diffusion barrier has a  
23 thickness of 100 to 1000 Å.
- 24 217. The EL laminate as set forth in claim 191, 202, 209 or 212, which further comprises,  
25 an injection layer above the dielectric layer, the second ceramic material or the barrier  
26 diffusion barrier, to provide a phosphor interface, composed of a binary, dielectric material  
27 which is non-stoichiometric in its composition and having electrons in a range of energy for  
28 injection into the phosphor layer.
- 29 218. The EL laminate as set forth in claim 217, wherein the injection layer is formed from a  
30 material which has greater than 0.5% atomic deviation from its stoichiometric composition.
- 31 219. The EL laminate as set forth in claim 218, wherein the injection layer is formed from  
32 hafnia or yttria.

1 220. The EL laminate as set forth in claim 219, wherein the injection layer has a thickness  
2 of 100 to 1000 Å.

3 221. The EL laminate as set forth in claim 218 or 220, wherein the injection layer is hafnia  
4 with a zinc sulfide phosphor, and wherein a diffusion barrier layer of zinc sulfide is used with  
5 a strontium sulfide phosphor.

6 222. A method of synthesizing strontium sulfide, comprising:

7 providing a source of high purity strontium carbonate in a dispersed form;

8 heating the strontium carbonate in a reactor with gradual heating up to a maximum  
9 temperature in the range of 800 to 1200°C;

10 contacting the heated strontium carbonate with a flow of sulfur vapours formed by  
11 heating elemental sulfur in the reactor to at least 300°C in an inert atmosphere; and

12 terminating the reaction by stopping the flow of sulfur at a point when sulfur dioxide or  
13 carbon dioxide in the reaction gas reaches an amount which correlates with an amount of  
14 oxygen in oxygen-containing strontium compounds in the reaction product which is in the  
15 range of 1 to 10 atomic percent.

16 223. The method as set forth in claim 222, wherein the sulfur is heated in the temperature  
17 range of 360 to 440°C.

18 224. The method as set forth in claim 222 or 223, wherein the strontium carbonate is  
19 provided in a dispersed form by mixing with one or more volatile, non-contaminating, clean  
20 evaporating compounds which decompose into gaseous products prior to the onset of the  
21 reaction of strontium carbonate.

22 225. The method as set forth in claim 224, wherein the volatile compound is selected from  
23 the group consisting of elemental sulfur and ammonium carbonate included in a weight ratio  
24 with strontium carbonate in the range of 1:9 to 1:1.

25 226. The method as set forth in claim 222 or 225, wherein the source of high purity  
26 strontium carbonate is doped with a source of cerium in the range of 0.01 to 0.35 mole%.

27 227. The method as set forth in claim 96, wherein the strontium sulfide phosphor is  
28 synthesized by a method comprising:

29 providing a source of high purity strontium carbonate in a dispersed form;

30 heating the strontium carbonate in a reactor with gradual heating up to a maximum  
31 temperature in the range of 800 to 1200°C;

32 contacting the heated strontium carbonate with a flow of sulfur vapours formed by

1 heating elemental sulfur in the reactor to at least 300°C in an inert atmosphere; and  
2 terminating the reaction by stopping the flow of sulfur at a point when sulfur dioxide or  
3 carbon dioxide in the reaction gas reaches an amount which correlates with an amount of  
4 oxygen in oxygen-containing strontium compounds in the reaction product which is in the  
5 range of 1 to 10 atomic percent.

6 228. The method as set forth in claim 227, wherein the sulfur is heated in the temperature  
7 range of 360 to 440°C.

8 229. The method as set forth in claim 227 or 228, wherein the strontium carbonate is  
9 provided in a dispersed form by mixing with one or more volatile, non-contaminating, clean  
10 evaporating compounds which decompose into gaseous products prior to the onset of the  
11 reaction of strontium carbonate.

12 230. The method as set forth in claim 229, wherein the volatile compound is selected from  
13 the group consisting of elemental sulfur and ammonium carbonate included in a weight ratio  
14 with strontium carbonate in the range of 1:9 to 1:1.

15 231. The method as set forth in claim 227 or 230, wherein the source of high purity  
16 strontium carbonate is doped with a source of cerium in the range of 0.01 to 0.35 mole%.

17 232. A method of forming a patterned phosphor structure having red, green and blue sub-  
18 pixel elements for an AC electroluminescent display, comprising:

19 a) selecting at least a first and a second phosphor, each emitting light in different  
20 ranges of the visible spectrum, but whose combined emission spectra contains red, green and  
21 blue light;

22 b) depositing a layer of the first phosphor which is to form at least one of the red, green  
23 or blue sub-pixel elements;

24 c) applying a photo-resist to the first phosphor, exposing the photo-resist through a  
25 photo-mask, developing the photo-resist, and removing the first phosphor in regions that the  
26 first phosphor is to define as one or more of the red, green and blue sub-pixel elements,  
27 leaving spaced first phosphor deposits, wherein the first phosphor is removed with an etchant  
28 solution comprising a mineral acid, or a source of anions of a mineral acid, in a non-aqueous,  
29 polar, organic solvent which solubilizes the reaction product of the first phosphor with anions  
30 of the mineral acid, and wherein optionally, prior to removing the first phosphor with the  
31 etchant solution, the first phosphor layer is immersed in the non-aqueous organic solvent;

32 d) depositing the second phosphor material over the first phosphor deposits and in

1 regions which are to define the other of the red, green and blue sub-pixel elements; and

2 e) removing by lift-off, the second phosphor material and the resist from above the first  
3 phosphor deposits leaving a plurality of repeating first and second phosphor deposits arranged  
4 in adjacent, repeating relationship to each other.

5 233. The method as set forth in claim 232, wherein the lift-off step is accomplished using a  
6 non-aqueous, predominately polar, aprotic solvent solution.

7 234. The method as set forth in claim 233, wherein at least one of the phosphors is an  
8 alkaline earth sulfide or selenide phosphor, and wherein the etchant solution is a mineral acid  
9 in methanol.

10 235. The method as set forth in claim 234, wherein the etchant solution includes an amount  
11 between 0.1 and 10% by volume of the mineral acid.

12 236. The method as set forth in claim 235, wherein the mineral acid is mineral acid is HCl  
13 or  $H_3PO_4$  or mixtures of these acids.

14 237. The method as set forth in claim 235 or 236, wherein the photoresist is a negative  
15 resist.

16 238. The method as set forth in claim 237, wherein the photoresist is a polyisoprene-based  
17 photoresist.

18 239. The method as set forth in claim 235, 237, or 238, wherein the lift-off is accomplished  
19 with a solution of methanol in toluene.

20 240. The method as set forth in claim 240, wherein the methanol is included in an amount  
21 between 5 and 20% by volume.

22 241. The method as set forth in claim 235, 237, 238 or 240, wherein one of the phosphors is  
23 a strontium sulfide phosphor.

24 242. The method as set forth in claim 241, wherein the first phosphor is a strontium sulfide  
25 phosphor, and the second phosphor is a zinc sulfide phosphor.

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